WO/2003/004591) METHOD FOR TRANSESTERIFICATION OF FATS AND/OR OILS BY MEANS OF ALCOHOLYSIS

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Applicants: PETER, Siegfried [DE/DE]; Lindenweg 3, 91080 Uttenreuth (DE).

WEIDNER, Eckhard [DE/DE]; Wasserstrasse 463a, 44795 Bochum (DE) (US Only).

Inventors: PETER, Siegfried; Lindenweg 3, 91080 Uttenreuth (DE).

WEIDNER, Eckhard; Wasserstrasse 463a, 44795 Bochum (DE).
BEYER, Andreas; Wuesthoff & Wuesthoff, Schweigerstrasse 2, 81541 München (DE).

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Title: (EN) METHOD FOR TRANSESTERIFICATION OF FATS AND/OR OILS BY MEANS OF

ALCOHOLYSIS

(DE) VERFAHREN ZUR UMESTERUNG VON FETT UND/ODER ÖL MITTELS ALKOHOLYSE

Abstract:

Agent:

(EN) The invention relates to a method for obtaining fatty acid esters, from triacyl glycerides by means of alcoholysis. The invention particularly relates to a method of the transesterification of fats and/or oils by alcoholysis, in which at least one alkyl fatty acid ester is added in such an amount during the initial phase that the resulting reaction mixture is a single phase. A higher reaction rate can thus be achieved with the method thereafter.

(DE) Die Erfindung betrifft ein Verfahren zur Gewinnung von Fettsäureestern aus Triacylglyceriden mittel Alkoholyse. Insbesondere betrifft die Erfindung ein Verfahren zur Umesterung von Fett und/oder Ol durch Alkoholyse, bei welchem zur Beschleunigung des Verfahrens in der Anfangsphase mindestens ein Alkanolfettsäureester in einer solchen Menge zugegeben wird, dass das dadurch entstehende Reaktionsgemisch einphasig wird. Dadurch Kann in dem Verfahren von vornherein eine hohe

Reaktionsgeschwindigkeit erhalten werden.

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Method to the transesterification of fat and/or oil by means of alcoholysis the invention relates to a method to the recovery of fatty add esters from Triacylylocretien by means of alcoholysis. In particular the invention concerns a method to the transesterification of fat and/or oil by alcoholysis, becomes added with which the acceleration of the method in the initial phase at least a Alkanollettsäureester in such an amount that the reaction mixture resultant thereby becomes single-bhase.

Transesterilication reactions actual are known. They represent a commercial significant class industrial organic reactions. With a transesterilication reaction an ester becomes by exchange of the acid groups or transers where the exchange of the alkoholschen groups into another ester. If the transesterilication becomes made by exchange of the alkoholschen groups, one speaks of the so called achorhysis (also Alkanolyse). With the alcoholysis the alcohol becomes and/or, the alkanols in the excess added, around an high yield at the desired ester to obtained. In ensert lime in connection with the generation of cliesel fluel from regenerating raw materials the preparation of < RTI ID=1.1> Alkyl star, </RTI> in particular of methyl resters, by vegetabilische oils < RTI ID=1.2> (z.</RTI> B. Rapsis). Solido: erheblich an Aktualität evennene.

The transesterification is an equilibrium reaction, which becomes usually already the triggered by mixing reactants. The reaction runs however so slow that is usually required for the commercial execution of the reaction a catalvist. As catalvist serve usually strong acidic ones or strong bases.

Fats and oils consist predominant of Glyceriden (mono, Di-und triglycerides).

With the transesterification of such lats and oils the component can become glycerol by low molecular monohydric alcohols substituted. In the practice for this the frequent method becomes after Bradshaw (described in the US patents < RTI ID=1.3 × 2, < IRTI > RTI ID=1.4 × 271, < IRTI> 619 and 2, 360, 844) applied. The reaction becomes conducted in an open container, which can consist of ordinary carbon steel. The lat or oil must be drying (anhydrous), clean and above all neutral, D. h. that content at free latty acids must be more negligible small (acid value) higher < as; RTI ID=1.5 × 1, < IRTI > 51.9 Generally the monohydric alcohol in high excess becomes added the increase of the yield and the reaction rate the reaction mixture (the equivalent ratio is often highers as IR.)

In a work of WRIGHTs et < RTI ID=2.3 at. / RTI > RTI ID=2.25 (Rtt. WRIGHT, J.c. / RTI> B. Segur, H. V. Clark, S. K. Coburn, E. E. Langdon and R. N. Durbis, oil & Soap, 2.1 [1944] 145-148) became the exact conditions for the alocholysis of lats with methanol and ethanol in the detail investigated. Further reported becomes from the authors over experiments over the abcoholysis with other monohydric alcohols. It becomes stated that the described above alcoholysis catalyzed with alkali is complete successful only if the fall is almost free of free fall yacids and the reaction mixture free of water. In one of these conditions is not mue, it comes to the saponification, which a loss to alkalinity and the formation of one; RTI ID=2.3> Gelstruktur / RTI> to the sequence, those has the separation and the setting of the dycerol prevented or slowed.

The transesterilication of the Triacylglyceride by means of alcoholysis is characterised in that the reaction between alkanois and Triacylglyceriden of a start-up phase requires, in which only a small reaction rate exists, because the reaction component is not alkanols in the oil soluble. Particularly with the preparation of methyl sesters this circumstance is very disturbing, since methanol is not he the unzuesternedno oils and lats. On the other hand methanol is in the methyl seters of the fatty acids good soluble. Pathways of the small < [ATI] ID = 2.4 Methanolic near that one. The Time of the transesterilication reaction truns of flow jowns one. The methanolic production of the Time of the transesterilication reaction truns of flow one. The mixture becomes single-phase and the reaction rate sudden strong rises.

As catalysts alkali motals or alcoholates of the alkali motals become used in the practice. The alkaline catalysts discover in the reaction mixture, D. h. the treaction becomes homogeneous catalyzed. The alkaline tacts and their alcoholates become in courses of the reaction soaps the reacted, which particularly separate in the resultant alcoholates become in courses of the reaction soaps the reacted, which particularly separate in the resultant playered rais whose subsequent retariment to pure dijector lates the price of. In addition, in the methyl ester are remain small amounts at alkali, which might not be problem-free whole with the use of the methyl ester as disestifued. Therefore also heteropenous catalyzed methods became, z in never time. B. using a metal sall of a strong basic amino acid as solid catalyst proposed (patent application DE 199 50 593 insoluble in the reaction mixture < RIII D=2.55 Aluminum, < / RII].

Furthermore a catalyst on basis of titanium oxides became developed, which it has the disadvantage that the reaction temperatures within the range of < RTI ID=3.1> 240 C< /RTI> lie.

On the basis of this state of the art it is to be eliminated the object of the invention this start-up phase with moderate reaction temperatures and/or, to shorten and to arrange the method thereby more effective.

The solution of this object made by a method to the transesterification of fat and/or oil by means of alcoholysis, with which the unzuesternden fat and/or oil the execution of the alcoholysis alkanois become, in particular univalent alkanois in the excess added, characterised in that the fat and/or the oil a least a Alkanoifettsäureester in such an amount added become that the reaction mixture resultant thereby becomes single-phase with reaction conditions.

Surprisingly it was now found that already a right small amount at supplied Alkanolfettsäureestern knows a this solve the problem. The addition of the Alkanolfettsäureester can take place at it forwards, after or simultaneous with the addition of the alkanols

After the method of this invention the initial phase becared he because it will also the the transesterilication thereby and/or.

After the method of this invention the initial phase becared to which se with a starting and and a starting and and a starting and and the starting and methyl estarting and methyl es

The fat and/or oil, which in the invention process used becomes, can be in particular biological origin.

The amount < on; RTI ID=3.9>. Alkanolitetsauresetern, < /RTI> to the preparation of a single-phase mixture added will must, hangs of the quality of the oil, which height < RTI ID=4.1>. Alkanolibberschusese</RTI> and the reaction temperature off. The Alkanolibberschuse becomes the increase of the reaction rate and the yield at Fettsauc RTI ID=4.2> realkanolestern</RTIS- generately with an equivalent ratio (D. h. Ratio of mol fatty acids in the fat and/or of 10 mol univalent alkanols) of 1: 6 or over it applied.

Alkanolfettsäureester, which in the method preferably added becomes, are z. B. Methyl ester, ethyl ester and/or Propylester.

< RTI ID=4.3> Alkanolfettsäureester
/RTI> preferably become in an amount from 5 to 50 Gew. - < %; RTI ID=4.4> particularly prefered 12 to 20 thread. - zugege < % related to the fat and/or oil; /RTI> user.

The invention process is particularly effective, if intended is, the transesterification in an heterogeneous catalyzed process, which preferably continuous can be to operate in addition, with an homogeneous catalyzed method the method is favourable after this invention, because the costs for the swirl of the two phases in the initial phase of the reaction saved to become to be able. Such heterogeneous catalyzed methods become for example in the mentioned above DF 199.65.054 described

Thus a catalyst becomes added in an other preferable embodiment the method, which can be either a soluble catalyst or in alkanols and in the reaction mixture insoluble metal salt of a amino acid or an amino acid derivative.

The solved catalyst can exist for example in solved alkali metals or Alkoholten of alkali metals.

The insoluble catayate an exhibit a metal component, which is calcium, strontium, barium, another alkaline and metal, or heavy metal, in particular silver, copper, zinc, manganes, to cobalt, lanthanum and on another rare earth metal, while the amino acid component of the insoluble catalyst can contain quarternaren nitrogen or a Countification of the Countification of the

The invention process can be accomplished to particularly effective, if the portion of free fatty acids in that < unzusesterndem fat and/or oil less than 0.5; RTI ID=5.1> Gew. - < %; /RTI> in particular less than 0.1 < RTI ID=5.2> Gew. - % < /RTI> amounts to.

In addition it was found that the reaction temperatures with the heterogeneous catalyzed transesterification preferably within the range of 80 to < RTI ID=5.3> 160 C, < /RTI> in particular within the range of 100 to < RTI ID=5.4> 150 C< /RTI> to lie should.

Particularly prefered becomes in the invention process a procedure, becomes recirculated with which the Alkanolletsfaureseter, which stays after separation of the glycerol from the product stream with the subsequent separation and purification of the major amount of the generated methyl esters by distillation as Sumplyrodukt. In this way simultaneous small amounts at not reacted Glycerolen become recirculated. In addition thereby one C RTI ID-5.5s. Glyceringehalt. / RTI-s in the final phase of the reaction reduced and the yield of the equilibrium reaction corresponding elevated. Altorether so a continuous processing becomes possible.

The amount at methyl esters, which < to the preparation of single-phaseness with reaction temperatures within the range of 100 to; RTI ID=5.6> 150 C< /RTI> prefered is, amounts to approx. 12 to 20 Gew. - %.

Subsequent one becomes the invention process at different examples near explained.

Thus the invention process at a mixture became from sunflower oil and methanol tested. In this case was sufficient with < RTI ID=5.7> 135 Cc < RTI> and an equivalent ratio of mol fatty acids in the oil to methanol of 1: 6 (60 Gew. < % sunflower oil and 40; RTI ID=5.8> Gew. < % < KTII> Methanol) an addition of approx. < RTI

|D=5.9 > 15% < /RII > RII |D=5.10 > Gew. "% < /RII > Methyl ester related to the oil, in order to produce an a phase system. The adjusting pressure amounted to in the described case <math>5 bar. & catalyst Z inkarigntal served. The Reaktionsgeschwin- RII |D=5.11> digkelit. /RII amounted to < RII |D=5.12> 2, 5 g/skgznag.</RII> In this example from the beginning an high reaction rate became obtained.

In addition palm oil was < with; RTI ID -5.13 - 150 C</RTI> with methanol in the equivalent ratio of 1: 6 mixed and Zinkarginia as catalyst added. After addition of 120 < RTI ID -5.14 > 60w. *Ve. /RTI> Methyl ester concerning palm oil was single-phase the mixture. The reaction rate was < with 3.2; RTI ID-6.1> g/skgznarg</RTI> very order in this initial phase with low reaction rate was immeed on.

Palm oil was < further with; RTI ID=6,2-8 5 C. /RTI> with methanol in the equivalent ratio of 1: 6 mixed and Zinkarginat as catalyst added. The reaction rate amounted to 0.05 < RTI ID=6.3-9 (skg.narg.; RTI> After by addition of methyl ester (approx. 13 < RTI ID=6.4-5 Gew. -% < /RTI> concerning oil the reaction mixture was sincle-shase. with ambient pressure a reaction rate of 0.0.3 % was < RTI ID=6.5 of \$6.8 capare. ATTI STIP measured.

With reaction temperatures of < RTI ID-6.65- 200</RTI- to < RTI ID-6.7> 240 C</RTI- become after methods with zinc soaps, described in the German patent specification DE 198 03 053 CI, as catalysts with printing up to 90 bar of triglycerides with high equivalent excess at methanol (equivalent ratio higher as 1:6) to seter seacted. Bottom these conditions is a higher methyl seter content emergency little, in order to manufacture an a phase system, when in the example mentioned with < RTI ID-6.65-135 C</RTI-

Claims < RTI ID= 7.1

- > 1.
 Int Method to the transesterification of fat and/or oil by means of alcoholysis, with which the unzuesternden fat and/or oil execution the aluminium becomes koholyse alkanols, in particular univalent alkanols in the excess added, characterised in that the fat and/or oil min destens < RIT ID=7.2> Alkanollettsauresster
 RIT in such an amount added becomes that the reaction mixture resultant thereby becomes sincle-phase.
- RTI ID=7.3> 2.
 /RTI> Process according to claim 1, characterised in that the admitted < RTI ID=7.4>
 Alkanolfettäureseter
 /RTI> selected becomes from the group, existing from ME thylestern, ethyl esters and/or Propylestern.
- 3. Process according to claim 1 or 2, characterised in that of the aluminium < RTI ID=7.5> kanolfettsäureester</ri>
 /RTI> in an amount from 5 to 50 < RTI ID=7.5> Gew. < %; /RTI> preferably 12 to 20 < RTI ID=7.7> Gew. < /ri>
 // RTI> referably 12 to 20 < RTI ID=7.7> Gew. </ri>
- 4. Process according to one of claims 1 to 3, characterised in that for the execution of the reaction in the reaction mixture of soluble Ka < RTI ID=7.8> talysator< /RTI> added becomes.
- 5. Process according to one of claims 1 < RTI ID=7.9> to 3, < /RTI> characterised in that for the execution of the reaction the reaction mixture in Alkano len and in the reaction mixture insoluble metal salt of a amino acid or an amino acid derivative an added become.
- Process according to claim 4, characterised in that the alcoholysis by solved alkali metals or alcoholates of the alkali metals catalyzed becomes.
- Process according to claim 5, characterised in that < RTIID=7.10> Metallkom < /RTI> ponente the catalyst calcium, strontium, barium, another earth alkene limetall or an heavy metal, in particular silver, copper, zinc, manganese,
- Iron, nickel, cobalt, lanthanum or another rare earth metal are.
- 8. Process according to one of claims 5 or 7, characterised in that the amino acid component of the catalyst quarternaren nitrogen or a Guanidinogruppe contains.
- 9. Process according to one of claims 5 or 7, characterised in that the catalyst a Schwermetallsalz of the arginine, in particular that
- Zinc salt or < RTI ID=8.1> Cadmiumsalz< /RTI> the arginine is.
- 10. Process according to one of claims 1 to 9, characterised in that the portion of free fatty acids in that unzuesterndem fat and/or oil < RTI ID=8.2> less than 0.5 thread. %, in particular less than 0.1 thread. btrâut. <%; //TII>
- < RTI |D=8.3> 11.< /RTI> Process according to one of claims 1 to < RTI |D=8.4> 10, < /RTI> characterised in that the transesterification with temperatures in the range from 80 to < RTI |ID=8.5> 160 C, < /RTI> preferably in the range from 100 to 150 C conducted becomes.
- 12. Process according to one of claims 1 to 11, characterised in that that the reaction mixture added < RTI 1D=8.65. Alkanollettsaureester < RTI> from the product stream into the method, resultant with the transesterification, rezirku one liert.

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